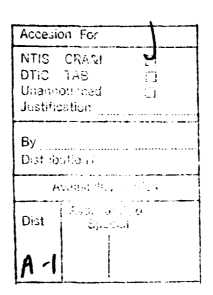
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Final discrepancy indices were R = 6.32% and wR = 4.97% for all 4947 independent data and R = 2.96% and wR = 3.67% for those 2924 reflections with  $|F_0| > 6\sigma(F_0)$ . The crystal is composed of K cations and  $(Me_3CCH_2)_3In-H-In(CH_2CMe_3)_3$  anions, each of which lie on two-fold axes. In-C(neopenty1) distances range from 2.199(6) to 2.231(6)Å (averaging 2.213Å), while In-( $\mu$ -H) distances are 1.933(15) and 1.950(23)Å. In-H-In angles are 151(5) and 161(5)°.





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Chemistry of Organoindium Hydrides. The Synthesis,

Characterization and Crystal Structure of K{H[In(CH<sub>2</sub>CMe<sub>3</sub>)<sub>3</sub>]<sub>2</sub>}

by

O. T. Beachley, Jr., Sun-Hua Chao, Melvyn Rowen Churchill and Ronald F. See

Prepared for Publication in Organometallics

State University of New York at Buffalo Department of Chemistry Buffalo, New York 14214

21 January 1992

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#### Chemistry of Organoindium Hydrides.

The Synthesis, Characterization and Crystal Structure of K{H[In(CH2CMe3)3]2}.

by

O. T. Beachley, Jr., \* Sun-Hua L. Chao, Melvyn Rowen Churchill \* and Ronald F. See

Department of Chemistry, State University of New York at Buffalo, Buffalo, NY 14214.

#### Abstract

Two new organoindium hydrides  $K\{H[In(CH_2CMe_3)_3]_2\}$  and  $K[HIn(CH_2CMe_3)_3]$ have been prepared and fully characterized according to their physical properties, partial elemental analyses, molecular weight studies and IR and  $^{1}\mathrm{H}$  and  $^{13}\mathrm{C}$  NMR spectral studies. The unique hydrogen atoms bonded to indium in the two compounds provide characteristic infrared bands and  $^{1}\mathrm{H}$  NMR resonances which have been identified by comparative studies with the corresponding deuterium derivatives. The indium hydride  $K\{H[In(CH_2CMe_3)_3]_2\}$ crystallizes in the monoclinic space group C2/c (No. 15) with cell parameters of  $\underline{a} = 22.243(5) \, \hat{A}$ ,  $\underline{b} = 17.021(3) \, \hat{A}$ ,  $\underline{c} = 21.290(3) \, \hat{A}$ ,  $\underline{B} = 17.021(3) \, \hat{A}$ 110.350(10)°, V = 7557(2)Å<sup>3</sup> and Z = 8. Final discrepancy indicies were R = 106.32% and wR = 4.97% for all 4947 independent data and R = 2.96% and wR = 3.67% for those 2924 reflections with  $|F_0| > 60(F_0)$ . The crystal is composed of  $K^+$  cations and  $(Me_3CCH_2)_3In-H-In(CH_2CMe_3)_3^-$  anions, each of which lie on two-fold axes. In-C(neopentyl) distances range from 2.199(6) to 2.231(6)Å (averaging 2.213Å), while In-( $\mu$ -H) distances are 1.933(15) and 1.950(23)Å. In-H-In angles are 151(5) and  $161(5)^{\circ}$ .

#### Introduction

Compounds which contain indium-hydrogen bonds are difficult to prepare, isolate and fully characterize. Thus, no indium hydride has been fully characterized previously by elemental analyses, molecular weight studies, diagnostic spectroscopic data and an X-ray structural study. The only compound with indium-hydrogen bonds for which an X-ray structural study has been reported is  $[(Me_3Si)_3C]In(H)(\mu-H)Li(THF)_2(\mu-H)In(\mu-H)(H)[C(SiMe_3)_3].$ 

However, the unique hydrogen atoms were not located. The presence of indium-hydrogen bonds was inferred from infrared and  $^6\text{Li}\{^1\text{H}\}$  NMR spectral data. This compound was too unstable to permit satisfactory carbon and hydrogen analyses. The closely related compounds M[InR<sub>3</sub>H] (M = Na, K; R = Me<sup>2</sup>, Et<sup>2</sup>, CH<sub>2</sub>SiMe<sup>3</sup><sub>3</sub>) are also unstable and decompose to MInR<sub>4</sub>, MH, indium metal and hydrogen. Thus, when In(CH<sub>2</sub>SiMe<sub>3</sub>)<sub>3</sub> and KH were combined in the absence of solvent, KIn(CH<sub>2</sub>SiMe<sub>3</sub>)<sub>4</sub> was isolated and was fully characterized.  $^3$ 

In this paper, the syntheses and characterizations of two novel compounds with indium hydrogen bonds,  $K\{H[In(CH_2CMe_3)_3]_2\}$  and  $K[HIn(CH_2CMe_3)_3]$ , are reported. Both compounds appear to be stable at room temperature and have been fully characterized.

#### Experimental

All compounds described in this investigation were extremely sensitive to oxygen and moisture and were manipulated in a standard vacuum line or under a purified argon atmosphere. The starting compound  $In(CH_2CMe_3)_3$  was prepared and purified by the literature method. 4 The reagents KH and KD were obtained from Aldrich Chemical Co. and were washed with pentane to remove oil prior to use. Solvents were dried by conventional procedures. Elemental analyses were performed by Schwarzkopf Microanalytical Laboratory, Woodside, NY or by E + R Microanalytical Laboratory, Inc., Corona, NY. Infrared spectra of Nujol mulls between CsI plates were recorded by means of a Perkin-Elmer 683 spectrometer. The <sup>1</sup>H NMR spectra were recorded at 400 MHz by means of a Varian VXR-400 S spectrometer, or at 300 MHz with a Varian Gemini-300 spectrometer. Proton chemical shifts are reported in  $\delta$  units (ppm) and are referenced to  ${\rm C_6H_6}$  at  $\delta$  7.15 ppm. The  $^{13}{\rm C\{}^1{\rm H\}}$  NMR spectra were recorded at 75 MHz by means of a Varian Gemini-300 spectrometer or at 101 MHz by means of a Varian VXR-400 S spectrometer. The proton-decoupled  $^{13}\text{C}$  spectra are reported relative to benzene at  $\delta$  128.0 ppm. All samples for NMR spectra were contained in sealed NMR tubes. Melting points were observed in sealed capillaries. Molecular weights were measured cryoscopically in benzene solution by using an instrument similar to that described by Shriver and Drezdzon.<sup>5</sup>

Synthesis of K[HIn(CH<sub>2</sub>CMe<sub>3</sub>)<sub>3</sub>]. The reagents  $In(CH_2CMe_3)_3$  (1.923 g, 5.858 mmol) and excess KH (0.583 g, 14.5 mmol) were combined in 20 mL pentane and the resulting mixture was stirred for 8 h at room temperature. The product was then separated from excess KH by extraction with pentane. Recrystallization of the crude product from pentane solution at -20 °C yielded K{HIn(CH<sub>2</sub>CMe<sub>3</sub>)<sub>3</sub>} (1.354 g, 3.676 mmol, 62.8% yield based on

In(CH<sub>2</sub>CMe<sub>3</sub>)<sub>3</sub>). Recrystallization at -20 °C of the concentrated mother liquor from the first recrystallization yielded additional K[HIn(CH<sub>2</sub>CMe<sub>3</sub>)<sub>3</sub>] (0.641 g, 1.74 mmol, 29.7% yield based on In(CH<sub>2</sub>CMe<sub>3</sub>)<sub>3</sub>). The two batches of crystals were identical. The total yield of purified K[HIn(CH<sub>2</sub>CMe<sub>3</sub>)<sub>3</sub>] was 92%.

Synthesis of K[DIn(CH<sub>2</sub>CMe<sub>3</sub>)<sub>3</sub>]. The deuteride K[DIn(CH<sub>2</sub>CMe<sub>3</sub>)<sub>3</sub>] was synthesized by using KD and the procedure previously desdribed. Recrystallizations from pentane yielded crystals of K[DIn(CH<sub>2</sub>CMe<sub>3</sub>)<sub>3</sub>].  $\frac{\text{K[DIn(CH<sub>2</sub>CMe<sub>3</sub>)<sub>3</sub>]}}{\text{K[DIn(CH<sub>2</sub>CMe<sub>3</sub>)<sub>3</sub>]}}. \quad \text{mp 117-120 °C decomp.} \quad ^{1}\text{H NMR (C<sub>6</sub>D<sub>6</sub>)} \quad \delta, \quad 0.63 \text{ (s, 6H, InCH<sub>2</sub>)}, \quad 1.29 \text{ (s, 27H, InCCCH<sub>3</sub>)}. \quad ^{13}\text{C{$^{1}$H}} \text{ NMR (C<sub>6</sub>D<sub>6</sub>)} \quad \delta, \quad 32.9 \text{ (CMe<sub>3</sub>)}, \quad 35.2 \text{ (CH<sub>3</sub>)}. \quad IR(Nujol mull, cm<sup>-1</sup>), \quad 1354 \text{ (s)}, \quad 1228 \text{ (s)}, \quad 1210 \text{ (s)}, \quad 1101 \text{ (m, sh)}, \quad 1076 \text{ (w)}, \quad ^{9}\text{90} \text{ (s, br spans region 1200-820)}, \quad 922 \text{ (w)}, \quad 904 \text{ (w)}, \quad 736 \text{ (m)}, \quad 699 \text{ (w, sh)}, \quad 659 \text{ (vs)}, \quad 649 \text{ (vs)}, \quad 561 \text{ (s, sh)}, \quad 448 \text{ (m, sh)}, \quad ^{405} \text{ (w, br spans region 500-300)}, \quad 372 \text{ (w)}, \quad 343 \text{ (m, sh)}, \quad 246 \text{ (m, sh)}, \quad 240 \text{ (m, sh)}.$ 

Synthesis of  $K\{H[In(CH_2CMe_3)_3]_2\}$ . Stoichiometric amounts of K[HIn(CH<sub>2</sub>CMe<sub>3</sub>)<sub>3</sub>] (0.431 g, 1.17 mmol) and In(CH<sub>2</sub>CMe<sub>3</sub>)<sub>3</sub> (0.385 g, 1.17 mmol) were combined in 30 mL of pentane. Initially, the mixture was a clear colorless solution, but after approximately 5 min a fine, colorless precipitate formed. The product was isolated after 10 min of stirring by cooling the mixture to -30 °C and filtering. Removal of the pentane left  $K\{H[In(CH_2CMe_3)_3]_2\}$  (0.793 g, 1.14 mmol, 97.2% yield). X-ray quality crystals were obtained by slow recrystallization of a pentane solution.  $K\{H[In(CH_2CMe_3)_3]_2\}$ . mp 137-143 °C decomp. Anal. Calcd. for  $C_{30}H_{67}InK$ : C, 51.73; H, 9.55. Found: C, 51.51; H, 9.60. <sup>1</sup>H NMR  $(C_6D_6)$   $\delta$ , 0.81 (s, 12H, InCH<sub>2</sub>), 1.25 (s, 54H, InCCCH<sub>3</sub>), 3.09 (br, 1H, InH).  $^{13}$ C{ $^{1}$ H} NMR (C<sub>6</sub>D<sub>6</sub>)  $\delta$ , 33.2 (CMe<sub>3</sub>), 35.4 (CH<sub>3</sub>). IR (Nujol mull, cm<sup>-1</sup>)  $\sim$ 1800-1100 (m, br), 1356 (s), 1348 (m), 1300 (vw, br), 1260 (w), 1228 (s), 1211 (m),  $\sim 1130-900$  (w, br), 1106 (m), 1092 (m), 1079 (w), 999 (w, sh), 981 (vw), 805 (w, br), ~760-630 (m, br), 734 (vw), 704 (vw), 679 (w), 645 (w), 563 (m), 449 (w), 373(w). Cryoscopic molecular weight, formula weight 696.60 (obsd molality, obsd mol wt, association): 0.0449, 729.9, 1.05; 0.0283, 746.4, 1.07; 0.0185, 783.0, 1.12.

Synthesis of  $K\{D[In(CH_2CMe_3)_3]_2\}$ . The deuteride  $K\{D[In(CH_2CMe_3)_3]_2\}$  was synthesized by using  $K[DIn(CH_2CMe_3)_3]$  and the procedure previously described.  $K\{D[In(CH_2CMe_3)_3]_2\}$ . mp 140-145 °C decomp. <sup>1</sup>H NMR ( $C_6D_6$ )  $\delta$ , 0.78 (s, 2H, InCH<sub>2</sub>), 1.26 (s, 9H, InCCCH<sub>3</sub>). <sup>13</sup> $C\{^1H\}$  NMR ( $C_6D_6$ )  $\delta$ , 32.8 (CMe<sub>3</sub>), 35.0 (CH<sub>3</sub>). IR (Nujol mull, cm<sup>-1</sup>), 1355 (s), 1349 (vw), 1229 (s), 1213 (s), 1103 (m), 1078 (w), ~1250-770 (s, br), 999 (m), 992 (m), 922 (w), 906 (w), ~750-600 (m, br), 735 (s), 700 (m), 661 (m), 650 (m), 600 (vw), 565 (vs, sh), ~500-350 (m, br), 470 (vw), 448 (m, sh), 406 (w), 374 (m), 343 (vw), 243 (m, sh).

Product Formed from In(CH2CMe3)3 and KH after 2 Hours,

K{H[In(CH<sub>2</sub>CMe<sub>3</sub>)<sub>3</sub>]<sub>2</sub>}. The reagents In(CH<sub>2</sub>CMe<sub>3</sub>)<sub>3</sub> (1.692 g, 5.153 mmol) and excess KH (0.703 g, 17.5 mmol) were combined in 20 mL pentane. After the resulting mixture was stirred for 2 h at room temperature, the crude product (0.547 g) was isolated by pentane extraction (x 6). The <sup>1</sup>H NMR spectrum identified the product as K{H[In(CH<sub>2</sub>CMe<sub>3</sub>)<sub>3</sub>]<sub>2</sub>}. <sup>1</sup>H NMR (C<sub>6</sub>D<sub>6</sub>)  $\delta$ , 0.78 (s, 2H, InCH<sub>2</sub>), 1.26 (s, 9H, InCCCH<sub>3</sub>), 3.05 (br, InH).

NMR Spectra of Mixtures of Components. The following experiments were performed in order to gain an understanding of the reactions occurring between KH,  $In(CH_2CMe_3)_3$ ,  $K[HIn(CH_2CMe_3)_3]$  and/or  $K\{H[In(CH_2CMe_3)_3]_2\}$ .

- A)  $In(CH_2CMe_3)_3$  and  $Excess\ K[HIn(CH_2CMe_3)_3]$ . Solvent (0.8 ml  $C_6D_6$ ) was added to a mixture of  $In(CH_2CMe_3)_3$  (0.0146 g, 0.0445 mmol) and excess  $K[HIn(CH_2CMe_3)_3]$  (0.0386 g, 0.105 mmol). <sup>1</sup>H NMR (8,  $C_6D_6$ ) 0.71 (s,  $InCH_2$ ), 1.28 (s,  $InCCCH_3$ ), 3.22 (s, InH).
- B)  $In(CH_2CMe_3)_3$  and  $Excess\ K\{H[In(CH_2CMe_3)_3]_2\}$ . Solvent (0.8 ml  $C_6D_6$ ) was added to a mixture of  $In(CH_2CMe_3)_3$  (0.0158 g, 0.0481 mmol) and excess  $K\{H[In(CH_2CMe_3)_3]_2\}$  (0.0450 g, 0.0646 mmol). <sup>1</sup>H NMR ( $\delta$ ,  $C_6D_6$ ) 0.85 (s,  $InCH_2$ ), 1.21 (s,  $InCCCH_3$ ), 2.97 (s, InH).
- C) In(CH<sub>2</sub>CMe<sub>3</sub>)<sub>3</sub> and Excess K[HIn(CH<sub>2</sub>CMe<sub>3</sub>)<sub>3</sub>]. The C<sub>6</sub>D<sub>6</sub> (0.8 ml) was added to a mixture of In(CH<sub>2</sub>CMe<sub>3</sub>)<sub>3</sub> (0.0123 g, 0.0375 mmol) and excess K[HIn(CH<sub>2</sub>CMe<sub>3</sub>)<sub>3</sub>] (0.0368 g, 0.0999 mmol). <sup>1</sup>H NMR ( $\delta$ , C<sub>6</sub>D<sub>6</sub>) 0.70 (s, InCH<sub>2</sub>), 1.28 (s, InCCCH<sub>3</sub>), 3.24 (s, InH). <sup>13</sup>C{<sup>1</sup>H} NMR (C<sub>6</sub>D<sub>6</sub>)  $\delta$ , 32.9 (CMe<sub>3</sub>), 35.1 (CH<sub>3</sub>).

Collection of the X-ray Diffraction Data for K{H[In(CH<sub>2</sub>CMe<sub>3</sub>)<sub>3</sub>]<sub>2</sub>}. A clear crystal having the dimensions  $0.25 \times 0.30 \times 0.40$  mm was aligned on a Siemens-upgraded Syntex P2<sub>1</sub>/R3 diffractometer. Data for ±h, k, ±l were collected by the coupled  $\theta$ (crystal)-2 $\theta$ (counter) scan technique and were

merged to a set of unique reflection. Details of data collection appear in Table I.

Axial photographs indicated that the crystal belonged to the C-centered subset of the monoclinic system. The systematic absences of hkl for h+k = 2n+1 and hOl for l=2n+1 indicate the possible space groups as Cc or C2/c. Intensity statistics clearly favored the centrosymmetric case. The space group C2/c  $[C_{2h}^6; No.15]$  was assumed and confirmed by the successful solution and refinement of the structure in this higher symmetry space group.

Solution and Refinement of the Crystal Structure. All crystallographic calculations were carried out on a VAX3100 workstation with the use of the Siemens SHELXTL PLUS program set. The analytical scattering factors for neutral atoms were corrected for the  $\Delta f'$  and  $i\Delta f''$  components of anomalous dispersion. The structure was solved by a combination of direct-methods and difference-Fourier techniques. Refinement led to convergence with R = 6.32%, wR = 4.97% and GOF = 0.95 for all 4947 reflections and 327 variables (R = 2.96% and wR = 3.67% for those 2924 reflections with  $F_O > 6.00(F_O)$ ).

All non-hydrogen atoms were refined anisotropically. The position and isotropic thermal parameter of the bridging hydride ions were refined. All hydrogen atoms of the neopentyl groups were included in calculated staggered positions with  $d(C-H) = 0.96 \text{\AA}^{-7}$ . A final difference-Fourier showed features cally in the range  $-0.61 + +0.83 \text{ e}^{-7}/\text{Å}^{-7}$ . Final atomic coordinates are collected in Table II.

#### Results and Discussion

Our studies of the reaction between  $In(CH_2CMe_3)_3$  and KH suggest that  $K[HIn(CH_2CMe_3)_3]$ , a simple Lewis acid-base adduct, is formed initially. However, this compound reacts rapidly with more  $In(CH_2CMe_3)_3$  to form

 $K\{H[In(CH_2,Me_3)_3]_2\}$ , the initial isolable product. If excess KH is available, then  $K\{H[In(CH_2CMe_3)_3]_2\}$  reacts to form  $K[HIn(CH_2CMe_3)_3]$  as the final product. The observation that  $K\{H[In(CH_2CMe_3)_3]_2\}$  can be isolated after  $In(CH_2CMe_3)_3$  and excess KH are stirred in pentane for two hours must be related to the limited solubility of  $K\{H[In(CH_2CMe_3)_3]_2\}$  and the insolubility of KH. When  $K[HIn(CH_2CMe_3)_3]$  and  $In(CH_2CMe_3)_3$  were combined in pentane, a solution was formed initially. However, after approximately 5 minutes, a fine, colorless precipitate of  $K\{H[In(CH_2CMe_3)_3]_2\}$  was observed and a quantitative yield was isolated after 10 min. A reaction mixture prepared from In(CH2CMe3)3 and excess KH must be stirred for 8 h in order to ensure the formation of  $K[HIn(CH_2CMe_3)_3]$ . Both new compounds have been characterized according to their physical properties, elemental analyses for carbon and hydrogen, cryoscopic molecular weight data, infrared spectra and  $^{1}\mathrm{H}$  and  $^{13}\mathrm{C}$  NMR spectra. The structure of  $\mathrm{K}\{\mathrm{H}[\mathrm{In}(\mathrm{CH}_{2}\mathrm{CMe}_{3})_{3}]_{2}\}$  in the solid state has also been elucidated by an X-ray structural study. Crystals of  $K[HIn(CH_2CMe_3)_3]$  were isolated also, but the diffraction data were too poor to be used in structure determination.

The crystal of  $K\{H[In(CH_2CMe_3)_3]_2\}$  is best viewed as an ordered array of  $K^+$  cations and  $(Me_3CCH_2)_3In-H-In(CH_2CMe_3)_3^-$  anions in a 1:1 ratio, all of which are bisected by a crystallographic two-fold axis. The crystallographic asymmetric unit consists of one half of two potassium cations (K(1) and K(2)) and one half of two  $[In(CH_2CMe_3)_3]_2H^-$  anions (centered about H(1) and H(2)). Figure 1 shows the mutual juxtaposition of two such crystallographic asymmetric units and the atomic labelling scheme. Interatomic distances (with esd's) are listed in Table III and interatomic angles are collected in Table IV.

The  $(Me_3CCH_2)_3In-H-In(CH_2CCMe_3)_3^-$  anions have precise  $C_2$  symmetry. The bridging hydride ligands are involved in 2-electron 3-center bonds. Observed indium-hydrogen distances are In(1)-H(1)=In(1A)-H(1)=1.950(23)Å, with (In(1)-H(1)-In(1A)=151(5)° and In(2)-H(2)=In(2A)-H(2)=1.933(15)Å with (In(2)-H(2)-In(2A)=161(5)°. The indium- $(\mu$ -hydride) distance is, as expected, substantially longer (by  $\sim 0.2$ Å) than the predicted terminal In-H distance of about 1.75Å (based on r(In)=1.44Å, from In-C(av)=2.21Å in the present structure less  $r(Csp^3)=0.77$ Å;  $^{8a}$  and r(H)=0.31Å from d(C-H)=1.08Å,  $^{8b}$  less  $r(Csp^3)=0.77$ Å).

The geometry about each indium atom is that of slightly flattened tetrahedron formed by the three  $\alpha$ -carbon atoms of the neopentyl groups and the bridging hydride ligand. The In-C bonds are equivalent within the limits of experimental error with In(1)-C(11) = 2.199(6), In(1)-C(21) =2.204(9) and In(1)-C(31) = 2.231(6)Å (aver = 2.211Å about In(1)) and In(2)-C(41) = 2.221(6), In(2)-C(51) = 2.200(7) and In(2)-C(61) = 2.222(6)Å (aver = 2.214Å about In(2)). These In-C distances in  $(Me_3CCH_2)_3In-H-In(CH_2CMe_3)_3^$ are similar to the corresponding distances in other organoindium anions. For example,  $KIn_2(CH_2SiMe_3)_{\mu}^3$  has an average In-C distance of 2.239[8]Å whereas that in LiIn[C(SiMe<sub>3</sub>)<sub>3</sub>]<sub>2</sub>H<sub>5</sub>•2THF is 2.20(3)Å; that in KInMe<sub>11</sub> is 2.239(3)Å; that in  $CsInMe_{\mu}^{9}$  is 2.26(2)Å and that in  $NaInPh_{\mu}^{10}$  is 2.230(3)Å. The In-C distances in these anionic species are longer than the In-C distances in most neutral, four coordinate indium compounds. Representative In-C distances in neutral species are 2.125(12) and 2.140(13)Å for  $[In(CH_2CMe_3)(CH_2SiMe_3)Cl]_2$ , 11 2.146(9) and 2.174(10)Å for  $[In(C_6Me_3H_2)_2Cl]_2^{12}$  and 2.182(6) and 2.210(7)Å for  $[(Me_3CCH_2)_2InPPh_2]_3$ . 13 One exception is the neutral organoindium compound [(Me<sub>3</sub>CCH<sub>2</sub>)<sub>2</sub>InCH<sub>2</sub>PPh<sub>2</sub>]<sub>2</sub>

whose In-C distances are comparable to those in  $K\{H[In(CH_2CMe_3)_3]_2\}$ ; the In-CH<sub>2</sub> distance is 2.258Å and the average In-C(neopenty1) distance is 2.222Å.

The C-In-C angles about the In(1) atom in  $K\{H[In(CH_2CMe_3)_3]_2\}$  are C(21)-In(1)-C(31)=114.9(3), C(11)-In(1)-C(31)=116.0(2) and  $C(11)-In(1)-C(21)=119.5(3)^\circ$  (aver = 116.8°). The C-In-C angles observed about In(2) are C(51)-In(2)-C(61)=115.8(3), C(41)-In(2)-C(61)=117.3(2) and  $C(41)-In(2)-C(51)=118.6(3)^\circ$  (aver = 117.2°). The C-In-H bond angles show values that are significantly smaller than the C-In-C angles. The values about In(1) are C(11)-In(1)-H(1)=104.3(24), C(21)-In(1)-H(1)=102.6(9) an  $C(31)-In(1)-H(1)=93.9(16)^\circ$  (aver = 100.3°). About In(2) the observed bond angles are C(41)-In(2)-H(2)=100.4(22), C(51)-In(2)-H(2)=101.4(8) and  $C(61)-In(2)-H(2)=97.1(15)^\circ$  (aver = 99.6°). This flattening of the tetrahedron, resulting in larger values for the C-In-C bond angles as compared to the C-In-H bond angles, is to be expected, since the neopentyl groups are substantially more bulky than the bridging hydride ligand (which is also electron deficient).

The potassium ions also lie on the crystallographic two-fold axis. They have their closest interactions with the hydride ligands  $(K(1) \cdot \cdot \cdot H(1) = 2.71(9) \text{Å}$  and  $K(2) \cdot \cdot \cdot \cdot H(2) = 2.87(8) \text{Å})$ . The  $K \cdot \cdot \cdot \text{In}$  distances are  $K(1) \cdot \cdot \cdot \cdot \text{In}(1) = K(1) \cdot \cdot \cdot \cdot \text{In}(1\text{A}) = 3.715(2) \text{Å}$  and  $K(2) \cdot \cdot \cdot \cdot \text{In}(2) = K(2) \cdot \cdot \cdot \cdot \text{In}(2\text{A}) = 3.721(2) \text{Å}$ . There are also some interactions between the potassium ions and hydrogen atoms on the neopentyl groups; these interactions are shown as dotted lines in Figure 1.

The two new indium hydrides are soluble in benzene and have been characterized by cryoscopic molecular weight data. The calculated molecular weights of the solutes in solutions prepared from  $K\{H[In(CH_2CMe_3)_3]_2\}$  reveal the presence of ion pairs with the simplest formula of the compound. In

contrast, solutions of  $K[HIn(CH_2CMe_3)_3]$  are composed of dimeric units,  $\{K[HIn(CH_2CMe_3)_3]\}_2$ . Because the calculated molecular weights of either compound do not vary significantly with concentration, neither associative nor dissociative equilibria are significant. Thus,  $K\{H[In(CH_2CMe_3)_3]_2\}$  can be envisioned to exist in solution as species related to the units identified in the solid state and to have a hydride ion serving as a Lewis base to two  $In(CH_2CMe_3)_3$  molecules as Lewis acids. Two alternative structures can be suggested for  $\{K[HIn(CH_2CMe_3)_3]\}_2$ . One structure involves five coordinate indium species with bridging hydride ions whereas the

$$Me_3CCH_2$$
 $Me_3CCH_2$ 
 $In -H$ 
 $K^+$ 
 $CH_2CMe_3$ 
 $Me_3CCH_2$ 
 $K^+$ 
 $CH_2CMe_3$ 
 $CH_2CMe_3$ 

alternative structure has tetrahedral  $HIn(CH_2CMe_3)_3$  anions with bridging potassium ions. The neopentyl group hydrogen atoms probably serve to coordinate with the potassium ions, as observed in the structural study. Available data do not permit us to distinguish between the two structures.

The  $^{1}$ H NMR spectra of K{H[In(CH<sub>2</sub>CMe<sub>3</sub>)<sub>3</sub>]<sub>2</sub>} and of K[HIn(CH<sub>2</sub>CMe<sub>3</sub>)<sub>3</sub>] have features which are directly related to hydride ions bonded to indium. The

<sup>1</sup>H NMR spectra of  $K[H[In(CH_2CMe_3)_3]_2]$  and  $K[HIn(CH_2CMe_3)_3]$  have broad lines at 3.09 and 3.41 ppm, respectively, which are not present in the spectra of the corresponding deuterium derivatives. Thus, these lines must be due to the unique hydrogen atoms bonded to indium. In contrast, the resonances of the methylene and methyl protons for the two indium hydride derivatives remain unchanged when the spectra of the respective protium and deuterium complexes are compared. It is also noteworthy that mixtures of  $K[H[In(CH_2CMe_3)_3]_2]$  and  $K[HIn(CH_2CMe_3)_3]$  exhibit only one set of resonances at chemical shifts which are intermediate between the two compounds. Thus, the neopentyl and hydride ligands exchange rapidly between the two compounds. Similarly, mixtures of  $K[H[In(CH_2CMe_3)_3]_2]$  and  $In(CH_2CMe_3)_3$  undergo rapid exchange of ligands as only one set of resonances is observed.

The infrared spectra of the two compounds are also consistent with the presence of indium hydrogen bonds. The compound  $K\{H[In(CH_2CMe_3)_3]_2\}$  has broad bands at ~1800-1100, ~1130-980 and ~760-630 cm<sup>-1</sup> which are partially obscured by Nujol and by other bands of the compound. These bands can be assigned to asymmetric stretching, symmetric stretching and bending modes for a In-H-In moiety which is bent rather than linear. These bands shift to ~1250-770, 750-600 and 500-350 cm<sup>-1</sup> upon deuterium substitution and are consistent with the results of reduced mass calculations. In comparison,  $K[HIn(CH_2CMe_3)_3]$  has broad, partially obscured bands at 1400-1000 and 900-500 cm<sup>-1</sup> which shift to 1200-820 and 500-300 cm<sup>-1</sup>, respectively, upon deuterium substitution. The bands centered about 1250 cm<sup>-1</sup> and 990 cm<sup>-1</sup> are assigned to In-H and In-D stretching modes whereas the lower frequency bands are due to bending modes. The shifts of the bands upon deuterium substitution is less than that expected by calculations based on reduced

masses. Thus, the bands in  $K[HIn(CH_2CMe_3)_3]$  probably are due to combinations of various modes and are not pure In-H vibrations.

The two new indium hydrides  $K\{H[In(CH_2CMe_3)_3]_2\}$  and  $K[HIn(CH_2CMe_3)_3]$  are inert to decomposition reactions in benzene solution over long periods of time. Neopentane  $CMe_4$  was not observed to be present in benzene solutions of the compounds, even after three months of standing at room temperature. In contrast, the closely related compound  $K[HIn(CH_2SiMe_3)_3]^3$  was observed to decompose in benzene solution to form indium metal,  $SiMe_4$ , biphenyl and varying amounts of  $H_2$ . All  $CH_2SiMe_3$  groups were converted to  $SiMe_4$  within 2 months at room temperature. These decomposition reactions of  $K[HIn(CH_2SiMe_3)_3]$  were proposed to be related to the formation of  $KCH_2SiMe_3$  by the dissociation of  $CH_2SiMe_3$ . Thus, the stability of the neopentyl derivatives suggest increased inertness to dissociative reactions, a property which might be related to the Lewis acidity of trineopentylindium. It is noteworthy that the reagents, 1 mmol of InMe3 and excess KH, formed 0.159 mmol of  $CH_4$  and indium metal in 12 h at 25 °C. 15

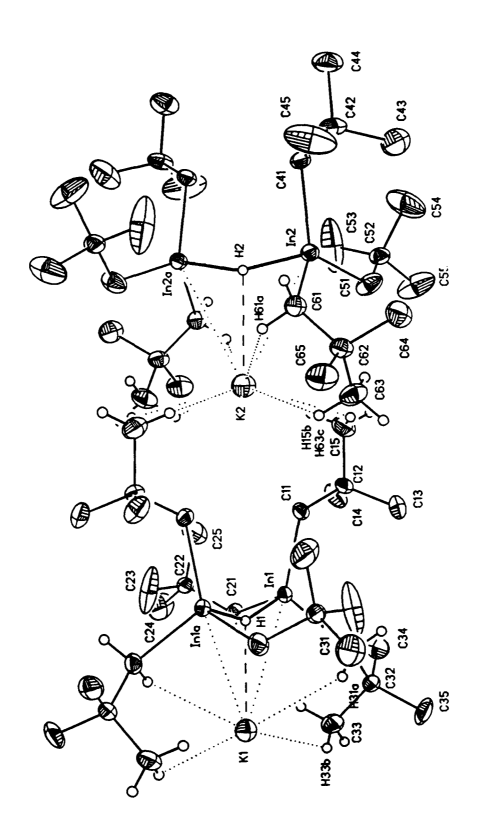
Acknowledgement. This work was supported in part by the Office of Naval Research and by a grant from Eastman Kodak Co. We thank the National Science Foundation for providing funds for the upgrade of the  $P2_1$  diffractometer to a Siemens  $P2_1/R3m$  through a grant from the Chemical Instrumentation Program (89-13733).

Supplementary Material. Anisotropic thermal parameters, calculated positions of hydrogen atoms and  $F_o/F_c$  list for  $K\{H[In(CH_2CMe_3)_3]_2\}$  (pages). For ordering information, see any current masthead page.

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Figure 1. The structure of  $K\{H[In(CH_2CMe_3)_3]_2\}$ . Hydrogen atoms of the neopentyl groups are omitted unless they interact with  $K^+$  ions. Note that a crystallographic two-fold axis passes through  $K(1) \bullet \bullet \bullet H(1) \bullet \bullet \bullet K(2) \bullet \bullet \bullet H(2)$ 



<u>Table I.</u> Experimental Data for the X-Ray Diffraction Study on  $K\{H[In(CH_2CMe_3)_3]_2\}$ .

# Crystal Data

Empirical Formula	$c_{30}^{}$ $H_{67}^{}$ $In_{2}^{}$ $K$
Color; Habit	clear crystal
Crystal Size (mm)	.40 x .30 x .25
Crystal System	Monoclinic
Space Group	C2/c
Unit Cell Dimensions	$\underline{a} = 22.243(5) \text{ Å}$
	b = 17.021(3)  Å
	$\underline{c} = 21.290(3) \text{ Å}$
	$\beta = 110.350(10)^{\circ}$
Volume	7557(2) Å <sup>3</sup>
2	8
Formula weight	696.6
Density(calc.)	$1.224 \text{ Mg/m}^3$
Absorption Coefficient	1.326 mm <sup>-1</sup>
F(000)	2912

### Data Collection

Diffractometer Used

Siemens R3m/V

Radiation

MoK $\alpha$  ( $\lambda = 0.71073 \text{ Å}$ )

Temperature (K)

295

Monochromator

Highly oriented graphite crystal

2θ Range

8.0 to 45.0°

Scan Type

 $2\theta - \theta$ 

Scan Speed

Constant;  $1.75^{\circ}/\text{min.}$  in  $\omega$ 

Scan Range  $(\omega)$ 

 $0.80^{\circ}$  plus K $\alpha$ -separation

Background Measurement

Stationary crystal and stationary counter at beginning and end of scan, each for 25.0% of total

scan time

Standard Reflections

3 measured every 97 reflections

Index Ranges

 $-24 \le h \le 24$ ,  $0 \le k \le 18$ 

 $-22 \le \ell \le 22$ 

Reflections Collected

10270

Independent Reflections

 $4947 (R_{int} = 2.74%)$ 

Observed Reflections

4947 (F >  $0.5\sigma(F)$ )

Absorption Correction

Semi-empirical

Min./Max. Transmission

0.3912 / 0.4473

## Solution and Refinement

System Used Siemens SHELXTL PLUS (VMS)

Solution Direct Methods

Refinement Method Full-Matrix Least-Squares

Quantity Minimized  $\sum w(F_0 - F_c)^2$ 

Absolute Structure N/A

Extinction Correction N/A

Hydrogen Atoms Riding model, fixed isotropic U

Weighting Scheme  $w^{-1} = \sigma^2(F) + 0.0008F^2$ 

Number of Parameters refined 327

Final R indices (obs. data) R = 6.32 %, wR = 4.97 %

R Indices (all data) R = 6.32 %, wR = 4.97 %

Goodness-of-Fit 0.95

Largest and Mean  $\Delta/\sigma$  0.003, 0.001

Data-to-Parameter Ratio 15.1:1

Largest Difference Peak 0.83 eÅ<sup>-3</sup>

Largest Difference Hole -0.61 eÅ<sup>-3</sup>

<u>Table II</u>. Atomic coordinates  $(x10^4)$  and equivalent isotropic displacement coefficients  $({\mathbb A}^2x10^3)$ .

	x	у	z	Ŭ(eq)
In(1)	-874(1)	11624(1)	1949(1)	40(1)
In(2)	222(1)	16625(1)	1712(1)	45(1)
K(1)	0	9745(2)	2500	83(1)
H(1)	0	11340(51)	2500	79(27)
K(2)	0	14748(2)	2500	80(1)
H(2)	0	16434(46)	2500	69(26)
C(11)	-818(3)	12884(3)	1747(3)	48(2)
C(12)	-1354(3)	13248(3)	1150(3)	53(3)
C(13)	-1320(5)	12929(5)	497(3)	97(4)
C(14)	-2014(3)	13061(5)	1181(4)	86(4)
C(15)	-1302(4)	14145(4)	1156(4)	90(4)
C(21)	-1433(4)	11233(5)	2567(3)	73(3)
C(22)	-1409(3)	11662(3)	3210(3)	52(3)
C(23)	-746(5)	11656(9)	3694(4)	181(8)
C(24)	-1834(6)	11263(6)	3537(5)	135(6)
C(25)	-1620(6)	12475(6)	3071(5)	152(8)
C(31)	-900(3)	10810(4)	1119(3)	54(3)
C(32)	-1465(3)	10240(4)	812(3)	53(2)
C(33)	-1458(3)	9590(4)	1302(4)	77(3)
C(34)	-2094(3)	10677(4)	642(4)	81(3)
C(35)	-1415(5)	9858(5)	174(4)	105(4)
C(41)	442(3)	17901(3)	1829(3)	53(3)
C(42)	882(4)	18253(4)	1501(4)	71(4)
C(43)	655(7)	18047(6)	771(6)	163(8)
C(44)	898(5)	19139(5)	1560(5)	109(5)
C(45)	1544(5)	17963(7)	1822(8)	174(9)
C(51)	-657(3)	16242(5)	910(4)	87(4)
C(52)	-1273(3)	16724(4)	695(3)	60(3)
C(53)	-1457(5)	16934(10)	1264(5)	213(10)
C(54)	-1160(5)	17476(7)	386(6)	168(7)
C(55)	-1814(4)	16315(6)	168(6)	160(6)
C(61)	1047(3)	15802(4)	1962(3)	58(3)
C(62)	1144(3)	15210(4)	1458(3)	61(3)
C(63)	620(4)	14592(4)	1267(4)	83(4)
C(64)	1105(4)	15653(5)	826(4)	98(5)
C(65)	1798(4)	14800(5)	1746(4)	96(4)

<sup>\*</sup> Equivalent isotropic U defined as one third of the trace of the orthogonalized  $\mathbf{U}_{ij}$  tensor

<u>Table III</u>. Interatomic Distances (Å) for  $K\{H[In(CH_2CMe_3)_3]_2\}$ .

•			
In(1)-K(1)	3.715 (2)	In(1)-C(11)	2.199 (6)
In(1)-C(21)	2.204 (9)	In(1)-C(31)	2.231 (6)
In(1)-H(1)	1.950 (23)	In(2)-K(2)	3.721 (2)
In(2)-C(41)	2.221 (6)	In(2)-C(51)	2.200 (7)
In(2)-C(61)	2.222 (6)	In(2)-H(2)	1.933 (15)
K(1)-H(31A)	2.757	K(1)-H(33B)	2.780
K(1)-H(1)	2.714 (87)	K(1)-In(1A)	3.715 (2)
K(1)-H(31C)	2.757 (6)	K(1)-H(33D)	2.780 (6)
K(1)-H(44D)	3.050 (11)	K(1)-H(44E)	3.050 (11)
K(1)-H(44F)	2.965 (11)	K(1)-H(44G)	2.965 (11)
K(2)-H(15B)	3.007	K(2)-H(15C)	3.024
K(2)-H(61A)	2.761	K(2)-H(63C)	2.751
K(2)-H(2)	2.869 (79)	K(2)-In(2A)	3.721 (2)
K(2)-H(15D)	3.007 (7)	K(2)-H(15E)	3.024 (7)
K(2)-H(61C)	2.761 (7)	K(2)-H(63D)	2.751 (10)
C(11)-C(12)	1.539 (7)	C(12)-C(13)	1.519 (11)
C(12)-C(14)	1.525 (11)	C(12)-C(15)	1.531 (9)
C(21) - C(22)	1.535 (10)	C(22)-C(23)	1.475 (10)
C(22) - C(24)	1.518 (15)	C(22)-C(25)	1.459 (12)
C(31) - C(32)	1.542 (8)	C(32)-C(33)	1.515 (10)
C(32)-C(34)	1.513 (9)	C(32)-C(35)	1.544 (11)
C(41) - C(42)	1.509 (12)	C(42)-C(43)	1.501 (14)
C(42)-C(44)	1,512 (10)	C(42)-C(45)	1.476 (13)
C(51)-C(52)	1,524 (10)	C(52)-C(53)	1.452 (14)
C(52)-C(54)	1.498 (14)	C(52)-C(55)	1.503 (11)
C(61)-C(62)	1.541 (10)	C(62)-C(63)	1.517 (10)
C(62)-C(64)	1.518 (11)	C(62)-C(65)	
H(1)-In(1A)	1.950 (23)	H(2)-In(2A)	1.933 (15)
( - / / /	. , - ,		

<u>Table IV</u>. Interatomic Angles (°) for  $K\{H[In(CH_2CMe_3)_3]_2\}$ .

K(1) - In(1) - C(11)	147.4(2)	K(1) - In(1) - C(21)	83.9(2)
C(11) - In(1) - C(21)	119.5(3)	K(1) - In(1) - C(31)	65.3(1)
C(11) - In(1) - C(31)	116.0(2)	C(21)-In(1)-C(31)	114.9(3)
K(1) - In(1) - H(1)	45.1(24)	C(11) - In(1) - H(1)	104.3(24)
C(21)-In(1)-H(1)	102.6(9)	C(31) - In(1) - H(1)	93.9(16)
K(2)-In(2)-C(41)	148.0(2)	K(2) - In(2) - C(51)	82.8(2)
C(41) - In(2) - C(51)	118.6(3)	K(2) - In(2) - C(61)	65.0(2)
C(41) - In(2) - C(61)	117.3(2)	C(51) - In(2) - C(61)	115.8(3)
K(2) - In(2) - H(2)	49.5(23)	C(41) - In(2) - H(2)	100.4(22)
C(51) - In(2) - H(2)	101.4(8)	C(61) - In(2) - H(2)	97.1(15)
In(1)-C(11)-C(12)	118.3(4)	C(11) - C(12) - C(13)	110.0(6)
C(11)-C(12)-C(14)	111.1(6)	C(13) - C(12) - C(14)	108.6(6)
C(11) - C(12) - C(15)	111.1(5)	C(13)-C(12)-C(15)	109.7(6)
C(14)-C(12)-C(15)	106.2(6)	In(1)-C(21)-C(22)	122.4(5)
C(21) - C(22) - C(23)	109.7(7)	C(21) - C(22) - C(24)	111.2(6)
C(23)-C(22)-C(24)	107.9(7)	C(21)-C(22)-C(25)	111.2(6)
C(23) - C(22) - C(25)	108.8(8)	C(24)-C(22)-C(25)	108.0(9)
In(1)-C(31)-C(32)	121.3(5)	C(31)-C(32)-C(33)	110.8(4)
C(31)-C(32)-C(34)	110.0(5)	C(33)-C(32)-C(34)	108.2(6)
C(31)-C(32)-C(35)	109.8(6)	C(33)-C(32)-C(35)	108.2(6)
C(34)-C(32)-C(35)	109.9(5)	In(2)-C(41)-C(42)	119.3(5)
C(41) - C(42) - C(43)	110.3(7)	C(41)-C(42)-C(44)	111.0(7)
C(43)-C(42)-C(44)	108.0(7)	C(41)-C(42)-C(45)	110.9(8)
C(43) - C(42) - C(45)	108.9(10)	C(44)-C(42)-C(45)	107.7(7)
In(2)-C(51)-C(52)	122.4(5)	C(51)-C(52)-C(53)	111.5(7)
C(51)-C(52)-C(54)	108.8(7)	C(53)-C(52)-C(54)	107.2(9)
C(51)-C(52)-C(55)	112.2(7)	C(53)-C(52)-C(55)	110.6(8)
C(54)-C(52)-C(55)	106.3(7)	In(2)-C(61)-C(62)	123.4(4)
C(61)-C(62)-C(63)	111.0(7)	C(61)-C(62)-C(64)	108.3(6)
C(63)-C(62)-C(64)	107.7(6)	C(61)-C(62)-C(65)	111.3(5)
C(63)-C(62)-C(65)	108.9(6)	C(64)-C(62)-C(65)	109.6(7)
In(1)-H(1)-K(1)	104.4(24)	In(1)-H(1)-In(1A)	151.3(49)
K(1)-H(1)-In(1A)	104.4(24)	In(2)-H(2)-K(2)	99.7(23)
In(2)-H(2)-In(2A)	160.6(46)	K(2)-H(2)-In(2A)	99.7(23)

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